

### THE UNITED STATES PATENT AND TRADEMARK OFFICE

Applicants:

Paul J. Coleman, et. al

Serial No.:

10/568,000

Case No. 21485YP

Art Unit:

1626

Filed:

February 10, 2006

Examiner:

For:

MITOTIC KINESIN INHIBITORS

Y.L. Chu

Commissioner for Patents P.O. Box 1450 Alexandria, VA 22313-1450

# DECLARATION OF CHRISTOPHER D. COX, PhD

Sir:

I Christopher D. Cox, PhD.,

# do hereby declare that

- I am employed by Merck & Co., Inc as a Senior Research Fellow in the department of Medicinal Chemistry. My curriculum vitae is attached as Exhibit
   1.
- 2. My responsibilities at Merck & Co., Inc. include basic oncology research. I worked extensively on the discovery of mitotic kinesin inhibitors. During the course of that work, my team and I discovered many mitotic kinesin inhibitors, including those claimed in the instant application.

3. The compounds of the instant invention are substituted pyrrolidines containing a fluorine in the 4-position. The addition of this fluorine in a strategic, metabolically benign location on the nitrogen containing ring unexpectedly results in a favorable profile when compared to unfluorinated analogues and analogues containing fluorines in other positions. The resulting compounds are more potent inhibitors of kinesin spindle protein (hereinafter "KSP"), and provide an acceptable Pgp ratio. This improvement can be demonstrated by comparing a fluorinated and unfluorinated analog:

Compound Structure	F N OH N N	F N OH
KSP IC <sub>50</sub>	12 nM	4.4 nM
Pgp ratio	34	3

The second compound is representative of those currently claimed in the instant application (See Scheme 2, page 74 in the specification).

4. These results were unexpected, and said results were realized prior to the filing date of the above-captioned application.

5. I hereby declare that all statements made herein of my own knowledge are true and that all statements made on information and belief are believed to be true; and further that these statements are made with the knowledge that willful false statements and the like so made are punishable by fine or imprisonment, or both under Section 1001 or Title 18 of the United States Code, and that such willful false statements may jeopardize the validity of the application of any patent issued thereon.

Christopher D. Cox, PhD

08-Dec-2008

Date

### Exhibit 1

#### **DECEMBER 2008**

### **CURRICULUM VITAE**

# I. PERSONAL

Christopher Cox, Ph. D. 422 Hoffman Road Harleysville, PA 19438 (267) 210-3474

# II. <u>EDUCATION</u>

Johns Hopkins University, Baltimore, MD
Organic Chemistry Ph. D.

Towson State University, Towson, MD
Chemistry
B. S. Summa Cum Laude

# III MERCK/MRL EMPLOYMENT HISTORY

Medicinal Chemistry

 Sr. Research Chemist
 11/2001 - 9/2004

 Research Fellow
 10/2004 - 3/2008

 Sr. Research Fellow
 4/2008 - Present

# IV. NON-MERCK EMPLOYMENT HISTORY

# V. ACADEMIC EXPERIENCE

NIH Postdoctoral Fellow Columbia University, New York, NY	1999 - 2001
Graduate Student Johns Hopkins University, Baltimore, MD	1994 - 1999
Undergrad. Res. Asst. Towson State U., Towson, MD	1994
NSF-REU Fellow University of Maryland, College Park, MD	1993

# VI. TRAINING

## VII. SOCIETY MEMBERSHIPS

Sigma Xi Member American Chemical Society

# VIII. ACADEMIC AND PROFESSIONAL HONORS

NIH Postdoctoral Fellowship	
Kilpatrick Graduate Fellowship	
ACS, Div. Of Organic Chem., Graduate Fellowship	
Ernest M. Marks Fellowship	
ACS, MD Section, Outstanding Student Award	1994
NSF-REU Fellowship	
Merck Index Achievement Award in Organic Chem.	

# IX. PUBLICATIONS AND PATENTS

#### **PUBLICATIONS**

- "Brazilian Baccharis Toxins: Livestock Poisoning and the Isolation of Macrocyclic Trichothecene Glucosides."
   B. Jarvis, S. Wang, <u>C. Cox</u>, M. Rao, V. Philip, M. Varaschin, C. Barros Natural Toxins 1996, 4, 58-71.
- "Copper(II)-Catalyzed Amide Isomerization: Evidence for N-Coordination."
   C. Cox, D. Ferraris, N. N. Murthy, T. Lectka
   J. Am. Chem. Soc. 1996, 118, 5332-5333
- "Crystal Structure and Triboluminescence 2. 9-Anthracenecarboxylic Acid and its Esters."
   L. Sweeting, A. Rheingold, J. Gingerich, A. Rutter, R. Spence, C. Cox, T. Kim Chem. Mater. 1997, 9, 1103-1115.
- 4. "Intramolecular Catalysis of Amide Isomerization." C. Cox, V. G. Young Jr., T. Lectka J. Am. Chem. Soc. 1997, 119, 2307-2308.
- 5. "Solvent Effects on the Barrier to Rotation in Carbamates." C. Cox, T. Lectka

  J. Org. Chem. 1998, 63, 2426-2427.
- 6. "Orthogonal" Lewis Acids: Catalyzed Ring Opening and Rearrangement of Acyl Aziridines."
  D. Ferraris, W. J. Drury III, C. Cox, T. Lectka
  J. Org. Chem. 1998, 63, 4568-4569

## **PUBLICATIONS** (continued)

7. "Intramolecular Catalysis of Amide Isomerization: Kinetic Consequences of the 5-NH--N<sub>a</sub> Interaction in Prolyl Peptides."

<u>C. Cox</u>, T. Lectka

J. Am. Chem. Soc. 1998, 120, 10660-10668.

8. "Diastereo- and Enantioselective Alkylation of α-Imino Esters with Enol Silanes catalyzed by R-Tol-BINAP-CuClO<sub>4</sub> (MeCN)<sub>2</sub>."
D. Ferraris, B. Young, <u>C. Cox</u>, W. J. Drury III, T. Dudding, T. Lectka *J. Org. Chem.* 1998, 63, 6090-6091.

9. "A Novel Synthesis of α-Amino Acid Derivatives through Catalytic Enantioselective Ene Réactions of α-Imino Esters."
W. J. Drury III, D. Ferraris, <u>C. Cox</u>, B. Young, T. Lectka J. Am. Chem. Soc. 1998, 120, 11006-11007.

"Strong Hydrogen Bonding to the Amide Nitrogen of an "Amide Proton Sponge": Consequences for Structure and Reactivity."
 C. Cox, H. Wack, T. Lectka
 Angew. Chem., Int. Ed. Engl. 1999, 38, 798-800.

11. "Nucleophilic Catalysis of Amide Isomerization." C. Cox, H. Wack, T. Lectka

J. Am. Chem. Soc. 1999, 121, 7963-7964.

12. "Intramolecular Acid-Catalyzed Amide Isomerization." <u>C. Cox</u>, T. Lectka *Org. Lett.* 1999, 1, 749-752.

13. "Synthetic Catalysis of Amide Isomerization." C. Cox, T. Lectka Acc. Chem. Res. 2000, 33, 849-858.

"Synthesis of the Functionalized Tricyclic Core of Lactonamycin by Oxidative Dearomatization."
 C. Cox, S. J. Danishefsky
 Org. Lett. 2000, 2, 3493-3496.

15. "Concise Synthesis of a Lactonamycin Model System by Diastereoselective Dihydroxylation of a Highly Fuctionalized Naphthoquinone."

<u>C. Cox</u>, S. Danishefsky

Org. Lett. 2001, 3, 2899-2902.

#### **PUBLICATIONS** (continued)

- 16. "Catalytic, Enatioselective Alkylation of α-Imino Esters: The Synthesis of Nonnatural α-Amino Acid Derivatives."
  - D. Ferraris, B. Young, <u>C. Cox</u>, T. Dudding, W. Drury III, L. Ryzhkov, A. Taggi, T. Lectka
  - J. Am. Chem. Soc. 2002, 124, 67-77.
- 17. "Studies Directed Toward the Total Synthesis of Lactonamycin: Control of the Sense of Cycloaddition of a Quinine Through Directed Intramolecular Catalysis."

  <u>C. Cox</u>, T. Siu, S. Danishefsky

  Angew. Chem., Int. Ed. Engl. 2003, 42, 5625-5629.
- 18. "Total Synthesis of Lactonamycinone."
  T. Siu, <u>C. Cox</u>, S. Danishefsky
  Angew. Chem., Int. Ed. Engl. 2003, 42, 5629-5634.
- "Two-Step Synthesis of β-Alkylchalcones and Their Use in the Synthesis of 3,5-Diaryl-5-Alkyl-4,5-Dihydropyrazoles."
   C. Cox, M. Breslin, B. Mariano
   Tetrahedron Lett. 2004, 45, 1489-1493.
- 20. "Kinesin Spindle Protein (KSP) Inhibitors. Part 1: The Discovery of 3,5-diaryl-4,5-dihydropyrazoles as Potent and Selective Inhibitors of the Mitotic Kinesin KSP."
  C. Cox; M.J. Breslin; B.J. Mariano; P.J. Coleman; C.A. Buser; E.S. Walsh; K. Hamilton; H.E. Huber; N.E. Kohl; M. Torrent; Y. Yan; L.C. Kuo; G.D. Hartman
  Bioorg. & Med. Chem. Lett. 2005, 15, 2041-2045.
- 21. "Kinesin Spindle Protein (KSP) Inhibitors. Part 4: Structure-based Design of 5-alkylamino-3,5-diaryl-4,5-dihydropyrazoles as Potent, Water-soluble Inhibitors of the Mitotic Kinesin KSP."
  C. Cox; M. Torrent, M.J. Breslin; B.J. Mariano; D.B. Whitman, P.J. Coleman; C.A. Buser; E.S. Walsh; K. Hamilton; M.D. Schaber, R.B. Lobell, W. Tao, V.J. South; N.E. Kohl; Y. Yan; L.C. Kuo; T. Prueksaritanont; D.E. Slaughter; C.Li; E. Mahan; B. Lu; G.D. Hartman Bioorg. & Med. Chem. Lett. 2006, 16, 3175-3179.

### **PUBLICATIONS** (continued)

- 22. "Kinesin Spindle Protein (KSP) Inhibitors. Part V: Discovery of 2-Propylamino-2,4-Diaryl-2,5-Dihydropyrroles as Potent, Water-Soluble KSP Inhibitors, and Modulation of their Basicity by β-Fluorination to Overcome Cellular Efflux by P-Glycoprotein."
  - C. Cox; M.J. Breslin; D.B. Whitman, P.J. Coleman; R.M. Garbaccio; M.E. Fraley; M.M. Zrada; C.A. Buser; E. S. Walsh; K. Hamilton; R. B. Lobell, W. Tao; M.T. Abrams; V.J. South; H.E. Huber; N.E. Kohl, G.D. Hartman. *Bioorg. & Med. Chem. Lett.* 17, 2697-2702 (2007).
- 23. "Kinesin spindle protein (KSP) inhibitors. Part 6: Design and synthesis of 3,5-diaryl-4,5-dihydropyrazole amides as potent inhibitors of the mitotic kinesin KSP."
  - Coleman, Paul J.; Schreier, John D.; Cox, Christopher D.; Fraley, Mark E.; Garbaccio, Robert M.; Buser, Carolyn A.; Walsh, Eileen S.; Hamilton, Kelly; Lobell, Robert B.; Rickert, Keith; Tao, Weikang; Diehl, Ronald E.; South, Vicki J.; Davide, Joseph P.; Kohl, Nancy E.; Yan, Youwei; Kuo, Lawrence; Prueksaritanont, Thomayant; Li, Chunze; Mahan, Elizabeth A.; Fernandez-Metzler, Carmen; Salata, Joseph J.; Hartman, George D. Bioorganic & Medicinal Chemistry Letters, 17(19), 5390-5395. (2007)
- "Kinesin Spindle Protein (KSP) Inhibitors. 9. Discovery of (2S)-4-(2,5-Difluorophenyl)-N-[3R,4S)-3-fluoro-1-methylpiperidin-4-yl)-N-methyl-2-phenyl-2,5-dihydro-1H-pyrrole-1-carboxamide (MK-0731) for the Treatment of Taxane-Refractory Cancer."
   C.D. Cox, P.J. Coleman, M.J. Breslin, D.B. Whitman, R.M. Garbaccio, M.E. Fraley...Y. Yan, ...G.D. Hartman

#### X. OTHER ACCOMPLISHMENTS

J. Med. Chem., 51, 4239-4252, (2008)

#### **INVITED LECTURES**

- "Discovery of L-001154704: A Potent and Selective Inhibitor of the Mitotic Kinesin KSP".
   Merck Chemistry Council Conference, La Sapiniere, Quebec - Canada August 2004
- "Intramolecular Catalysis of Amide Isomerization and its Role in Protein Folding".
   Towson State University, Towson, MD May 1998.

## **INVITED LECTURES (continued)**

 "Discovery of Kinesin Spindle Protein Inhibitor MK-0731 for the Treatment of Taxane-Refractory Cancer".
 Johns Hopkins University April 30, 2008:

## **PRESENTATIONS**

- "Discovery and optimization of kinesin spindle protein (KSP) inhibitors."
   <u>Cox, Christopher D.</u>; Coleman, Paul J.; Fraley, Mark E.; Garbaccio, Robert M.;
   Breslin, Michael J.; Whitman, David B.; Schreier, John D.; Hartman, George D.;
   Torrent, Maricel; Lobell, Rob; Buser, Carolyn; Tao, Weikang; Huber, Hans;
   Kohl, Nancy E.; Yan, Youwei; Kuo, Lawrence C.
   Abstracts of Papers, 233rd ACS National Meeting, Chicago, IL, United States,
   March 25-29, 2007 (2007), MEDI-211.
- "HTS to MK-0731: The Role of Fluorine in Optimization of Kinesin Spindle Protein (KSP) Inhibitors for the Treatment of Cancer." Spring ACS National Meeting, April 7, 2008 (2008)
- "Chemical Strategies to Alter P-Glycoprotein Efflux of Drug Molecules" *Spring ACS National Meeting*, April 8, 2008 (2008)